

Siemens Medical Solutions, USA, Inc. % Ms. Christine Dunbar Senior Regulatory Affairs Specialist 685 East Middlefield Road MOUNTAIN VIEW CA 94043 April 22, 2020

Re: K200585

Trade/Device Name: ACUSON SC2000 Diagnostic Ultrasound System

Regulation Number: 21 CFR 892.1550

Regulation Name: Ultrasonic pulsed doppler imaging system

Regulatory Class: Class II

Product Code: IYN, IYO, ITX, OBJ, LLZ

Dated: March 4, 2020 Received: March 6, 2020

Dear Ms. Dunbar:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see

https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to https://www.fda.gov/medical-device-problems.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance) and CDRH Learn (https://www.fda.gov/training-and-continuing-education/cdrh-learn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

For

Thalia T. Mills, Ph.D.
Director
Division of Radiological Health
OHT7: Office of In Vitro Diagnostics
and Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2020

See PRA Statement below.

510(k) Number (if known) K200585

Device Name

ACUSON SC2000 Diagnostic Ultrasound System

Indications for Use (Describe)

The SC2000 ultrasound imaging system is intended for the following applications: Cardiac, Neo-natal and Fetal Cardiac, Pediatric, Transesophageal, Adult Cephalic, Peripheral Vessel, Abdominal, Intraoperative Abdominal, Musculo-skeletal Conventional, and Musculo-skeletal Superficial applications. The system also provides the ability to measure anatomical structures and calculation packages that provide information to the clinician that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

The typical examinations performed using the SC2000 Ultrasound System are:

Cardiac Imaging Applications and Analysis

The system transmits ultrasound energy into adult, pediatric, neonatal, and fetal cardiac patients creating 2D (B), 3D, MMode (M), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave (PW) Doppler, and Continuous Wave Doppler (CWD) to obtain images and blood flow velocity of the heart, cardiac valves, great vessels, and surrounding anatomical structures to evaluate the presence or absence of pathology. The system may be used to acquire patient electrocardiogram for synchronizing the diastolic and systolic capture of ultrasound images.

The system also supports catheters which are intended for intra-cardiac and intraluminal visualization of cardiac and great vessel anatomy and physiology as well as visualization of other devices in the heart of adult and pediatric patients. The catheter is intended for imaging guidance only, not treatment delivery, during cardiac interventional percutaneous procedures. The system has Cardiac Measurements and Calculation Packages that provide information that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

Vascular Imaging Applications and Analysis

The system transmits ultrasound energy into various parts of the body of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave Doppler (PWD), and Continuous Wave Doppler (CWD) to obtain images and blood flow velocity of the carotid arteries or juggler veins in the neck; superficial and deep veins and arteries in the arms and legs and abdomen; and surrounding anatomical structures to evaluate the presence or absence of pathology. The system may be used to acquire patient electrocardiogram for synchronizing the diastolic and systolic capture of ultrasound images. The system has Vascular Measurements and Calculation Packages that provide information that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

Superficial Imaging Applications

The system transmits ultrasound energy into various parts of the body of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave Doppler (PWD), and Continuous Wave Doppler (CWD) to obtain images and blood flow velocity of conventional or superficial musculoskeletal structures and surrounding anatomical structures to evaluate the presence or absence of pathology. The system may be used to acquire patient electrocardiogram for synchronizing the diastolic and systolic capture of ultrasound images.

Intraoperative Imaging Applications

The system transmits ultrasound energy into various parts of the body of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), and Pulsed Wave Doppler (PWD) to obtain images and blood flow velocity that provide guidance during intraoperative procedures.

Transcranial Imaging Applications

The system transmits ultrasound energy into the cranium of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave Doppler (PWD), and Continuous Wave Doppler (CWD) to obtain images and blood flow velocity of the brain and surrounding anatomical structures to evaluate the presence or absence of pathology. The system provides Measurement Packages that provide information that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

Type of Use (Select one or both, as applicable)		
Prescription Use (Part 21 CFR 801 Subpart D)	Over-The-Counter Use (21 CFR 801 Subpart C)	

CONTINUE ON A SEPARATE PAGE IF NEEDED.

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SECTION 1. 2. 510(K) SUMMARY

Date: March 04, 2020 K200585

Part 1. Sponsor: Siemens Medical Solutions USA, Inc.,

Ultrasound Division

685 East Middlefield Road

Mountain View, California 94043

Contact Person: Christine Dunbar

Tel: (925) 374-2045

Part 2. Device Name: ACUSON SC2000 Diagnostic Ultrasound System

Common Name: Diagnostic Ultrasound System with Accessories

Classification: Regulatory Class: II

Review Category: Tier II

Classification Panel: 90, Radiology

Ultrasonic Pulsed Doppler Imaging 892.1550 90-IYN System Ultrasonic Pulsed Echo Imaging System 892.1560 90-IYO Diagnostic Ultrasound Transducer 892.1570 90-ITX Diagnostic Intravascular Catheter 892.1200 90-OBJ Picture archiving and Communications 892.2050 90-LLZ

System (Optional PACS for SC2000)

Manufacturing Site: 2500 Millbrook Drive, Suite B Buffalo Grove, Illinois USA

Legal Manufacturer: Siemens Medical Solutions USA, Inc.

685 East Middlefield Road Mountain View, CA 94043

Part 3. Legally Marketed Predicate Devices

The ACUSON SC2000 Diagnostic Ultrasound System is a multi-purpose diagnostic ultrasound system with accessories and proprietary software, and is substantially equivalent to the company's own products, the ACUSON SC2000, v5.1 (VB21) under K181098 on May 31, 2018, is the primary predicate device.

Part 4. Device Description

The ACUSON SC2000 Diagnostic Ultrasound System is a multi-purpose mobile, software controlled, diagnostic ultrasound system with an on-screen display of thermal and mechanical indices related to potential bio-effect mechanisms. Its function is to transmit and receive ultrasound echo data and display it in B-Mode, M-Mode, Pulsed (PW) Doppler Mode, Continuous (CW) Doppler Mode, Color Doppler Mode, Color M Mode, Doppler Tissue Mode, Amplitude Doppler Mode, a combination of modes and Harmonic Imaging on a Display. The transducer and catheter-based transducers will follow Track 3 acoustic labeling (AIUM 1004, IEC 2007, AIUM/NEMA 2004a).

Part 5. Intended Use and Indications for Use Statements

SC2000 Diagnostic Ultrasound System, VC10A (v6.0)

(The Indications for Use remains unchanged as cleared under K181098 (VB21A)

The SC2000 ultrasound imaging system is intended for the following applications: Cardiac, Neo-natal and Fetal Cardiac, Pediatric, Transesophageal, Adult Cephalic, Peripheral Vessel, Abdominal, Intraoperative Abdominal, Musculo-skeletal Conventional, and Musculo-skeletal Superficial applications. The system also provides the ability to measure anatomical structures and calculation packages that provide information to the clinician that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

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The system also supports catheters which are intended for intra-cardiac and intraluminal visualization of cardiac and great vessel anatomy and physiology as well as visualization of other devices in the heart of adult and pediatric patients.

The system has Cardiac Measurements and Calculation Packages that provide information that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

Vascular Imaging Applications and Analysis

The system transmits ultrasound energy into various parts of the body of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave Doppler (PWD), and Continuous Wave Doppler (CWD) to obtain images and blood flow velocity of the carotid arteries or juggler veins in the neck; superficial and deep veins and arteries in the arms and legs and abdomen; and surrounding anatomical structures to evaluate the presence or absence of pathology. The system may be used to acquire patient electrocardiogram for synchronizing the diastolic and systolic capture of ultrasound images.

The system has Vascular Measurements and Calculation Packages that provide information that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

Superficial Imaging Applications

The system transmits ultrasound energy into various parts of the body of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave Doppler (PWD), and Continuous Wave

Doppler (CWD) to obtain images and blood flow velocity of conventional or superficial musculoskeletal structures and surrounding anatomical structures to evaluate the presence or absence of pathology. The system may be used to acquire patient electrocardiogram for synchronizing the diastolic and systolic capture of ultrasound images.

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The system transmits ultrasound energy into various parts of the body of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), and Pulsed Wave Doppler (PWD) to obtain images and blood flow velocity that provide guidance during intraoperative procedures.

Transcranial Imaging Applications

The system transmits ultrasound energy into the cranium of adult patients creating 2D (B), Color Doppler (CD), Color Power Doppler (CPD), Pulsed Wave Doppler (PWD), and Continuous Wave Doppler (CWD) to obtain images and blood flow velocity of the brain and surrounding anatomical structures to evaluate the presence or absence of pathology.

The system provides Measurement Packages that provide information that may be used adjunctively with other medical data obtained by a physician for clinical diagnosis purposes.

Part 6. Summary of Technological Characteristics

6.1 The ACUSON SC2000 VC10A (v6.0) Diagnostic Ultrasound System is a multi-purpose diagnostic ultrasound system with accessories and proprietary software, and is substantially equivalent to the company's own products, the ACUSON SC2000 VB21A (v5.1) (K181098) with regard to both intended use and technological characteristics. Both the subject ultrasound system and the predicate ultrasound system function in the same manner as similar diagnostic ultrasound systems and transducers including the cardiac catheters such as the ACUSON V and Volume ICE catheters and support for third party ICE catheters; the SoundStarTM 8F and 10F and the SoundStarTM *eco* 8F and *eco* 10F versions of the Diagnostic Ultrasound Catheters, which are developed, manufactured and distributed by Biosense Webster, Inc. under multiple clearances.

The foundation of the ACUSON SC2000 VC10A (6.0) is the SC2000 VB21A (5.1) release with features and transducers integrated with the ACUSON SC2000 hardware and the software developed for SC2000 platform. The updated SC2000 system software VC10A includes the same currently cleared software applications and support for the optional SC2000 (WP) workplace.

It is Siemens' opinion that the ACUSON SC2000 VC10A (v6.0) is substantially equivalent to the predicate device, the ACUSON SC2000 VB21A (V5.1) with regard to both intended use, indications for use and technological characteristics.

6.2 - List of Technological Characteristics and SE Comparison Table

Feature / Characteristic	ACUSON SC2000 v6.0 Including Transducers & Catheters This Submission	ACUSON SC2000 5.1 K181098 Including Transducers & AcuNav Volume ICE Catheter & 3 rd Party Catheters.
Indications for Use:		
Indications for Use - Device	Unchanged for VC10A (v6.0)	V

Feature / Characteristic	ACUSON SC2000 v6.0 Including Transducers & Catheters This Submission	ACUSON SC2000 5.1 K181098 Including Transducers & AcuNav Volume ICE Catheter & 3 rd Party Catheters.
■ Fetal	$\sqrt{}$	√ V
Abdominal	V	V
Pediatric	V	V
■ Small Organ	$\sqrt{}$	√
■ Cardiac	√	√
Peripheral vessel	$\sqrt{}$	$\sqrt{}$
Musculo-skeletal (conventional)	$\sqrt{}$	$\sqrt{}$
Musculo-skeletal (superficial)	V	V
Frequencies Supported:	$\sqrt{\frac{1.7\text{MHz}}{10\text{MHz}}}$	(1.7MHz~10MHz)
Modes:		
■ B	V	V
■ M	V	V
PWD (Pulsed Wave Doppler)	V	√
 CWD (Continuous Wave Doppler) 	$\sqrt{}$	$\sqrt{}$
 PW DTI (Doppler Tissue Image) 	$\sqrt{}$	V
Color Doppler	$\sqrt{}$	$\sqrt{}$
Color Power Doppler (CPD)	$\sqrt{}$	$\sqrt{}$
Combined (BMDC)	\checkmark	$\sqrt{}$
Transducers/ Catheters		
Z6Ms - TEE Transducer	V	√
4Z1c - Phased Array Transducer	$\sqrt{}$	V
	optimized factory default preset for 4V1c	Preset available.
10V4 - Phased Array Transducer	V	V
9L4 - Linear Array Transducer	$\sqrt{}$	$\sqrt{}$
9L4 - Linear Array Transducer – update imaging presat	optimized factory default preset for 9L4	Preset available.
4V1c - Phased Array Transducer	V	V
V5Ms - TEE Transducer	· √	· √
8V3 - Phased Array Transducer	, √	√
	√ √	√ √
V7M - TEE Transducer	V	v a/
6C1HD - Curved array transducer CW2 - Continuous Wave Transducer	√ √	\ √

Feature / Characteristic	ACUSON SC2000 v6.0 Including Transducers & Catheters This Submission	ACUSON SC2000 5.1 K181098 Including Transducers & AcuNav Volume ICE Catheter &
AcuNav 8F - Phased Array Ultrasound Catheter	V	3 rd Party Catheters. √
AcuNav 10F Intracardiac Transducer	$\sqrt{}$	V
AcuNav™ V ICE Catheter	V	V
AcuNav™ Volume ICE Catheter (aka P6)	V	V
AcuNav [™] Volume ICE Catheter Imaging Preset update	optimized factory default preset for AcuNav Volume	Preset available.
Supports Third party US Catheters with Swift-Link connector:	1	1
SoundStar [™] eco 8 French catheter ¹	V	V
SoundStar [™] <i>eco</i> 10 French catheter	V	V
SoundStar [™] 8 French catheter ²	V	V
SoundStar™ 10 French catheter	V	V
SwiftLink Connector	V	V
Features on SC2000 System:		
Patient Registration Fields	Improve Patient data entry workflow	V
 Change/Edit Patient Information Active exam 	Improve Patient data entry workflow	Edit non-active exam only
 Native[™] tissue harmonic imaging (2D Brightness mode) 	V	(Harmonic imaging)
■ TEQ™ ultrasound technology	V	(Updated TEQ)
■ Volume ICE Package	V	V
 Support for AcuNav Volume ICE Catheter connected by SwiftLink cable. 	√	AcuNav Volume ICE Catheter connected by SwiftLink Catheter supported.

 $^{^{1}}$ SoundStar and SoundStar eco catheters are products of Biosense Webster, Inc. (A Johnson & Johnson company) holds the 510(k) clearances for these devices.

² SoundStar and SoundStar eco catheters are products of Biosense Webster, Inc. (A Johnson & Johnson company) holds the 510(k) clearances for these devices.

Feature / Characteristic	ACUSON SC2000 v6.0 Including Transducers & Catheters This Submission	ACUSON SC2000 5.1 K181098 Including Transducers & AcuNav Volume ICE Catheter & 3 rd Party Catheters.
 ICE Catheter Auto- reduce Mechanical Index (MI) when imaging in air in order to protect the catheter from thermal damage and preserve image quality. 	New feature for V & Volume ICE and Sound Star ICE catheters.	-
■ 2D ICE Package	(Unchanged)	√
 True Volume Imaging Support, AcuNav Volume ICE Catheter 90° x 50° real-time volume imaging 	√ (Unchanged)	V
■ Volume Color Doppler	V	V
Fetal Echo Calculations	V	V
■ Fetal Imaging Presets	V	V
 Cardiac Imaging physiological signal display 	V	V
 eSie Measure on TEE 	Update Trace	V
■ Advanced SieClear [™] spatial compounding	V	V
 Clarify™ vascular enhancement technology 	V	\checkmark
syngo ® Velocity Vector Imaging (eSie VVI)	Enhancement – Bullseye red/blue color map	V
 eSie Valves Advanced Measurement Package 	V	V
■ eSie LVA	V	V
eSie LVA: 5 Cardiac Cycles	Update to number of cardiac cycles supported.	Currently 3 cardiac cycles
■ eSie PISA	V	V
 eSie Left Heart Measurement Package 	V	V
Volume Right Ventricular Analysis (RVA)	√	V
Stress Echo Package	$\sqrt{}$	$\sqrt{}$
 Rapid Stress Volume Stress Echo App. 	√	√
 Cardiac Measurements and Calculations 	√	√

Feature / Characteristic	ACUSON SC2000 v6.0 Including Transducers & Catheters This Submission	ACUSON SC2000 5.1 K181098 Including Transducers & AcuNav Volume ICE Catheter & 3 rd Party Catheters.
syngo ® TrueFusion v1.0	V	V
Septal Guide – Orthogonal Guidelines	V	V
VR Measurement Tools (Volume)	The ability to measure anatomy and pathology directly on the Volume Rendered (VR) images in CINE and acquired in 4D.	_
Reference Lines One-click MPR alignment	New Display Feature	-
Volume Reference Line Projections eSie Slice / eSie Lines	New Display Feature	_
One-click MPR A/B Align on Volume Review (VR)	New Display Feature	-
Trace erase, back up behavior with Trackball: B- Mode	Updated Display Feature	V
D'Art renamed Single V	Consolidated windows	√
Contrast Agent Imaging	\checkmark	V
 Edit patient data on active exam. 	Edit patient data on active exam	Edit patient data on completed exam
■ Zoom & Pan	$\sqrt{}$	$\sqrt{}$
eSieScan (Protocols)	\checkmark	$\sqrt{}$
InFocus Coherent Technology	\checkmark	V
Circle Tool	V	V
■ DICOM		√ √
DICOM SR (Structure Reports)	V	V
 DICOM GSDF Monitor Support 	DICOM GSDF (Gray scale Standard Display Function) review monitor to look comparable to those displayed on the SC2000 system monitor.	-

	ACUSON SC2000 v6.0 Including	ACUSON SC2000 5.1 K181098 Including Transducers
Feature / Characteristic	Transducers & Catheters	&
	This Submission	AcuNav Volume ICE Catheter & 3 rd Party Catheters.
■ DICOM Tags (Teamplay)	DICOM support the export of DICOM tags for System Data Collection (SDC) / Utilization	-
MS Windows 10 migration	Management (UM) Windows 10	Windows 7
Wireless - enabled	VIIIdows 10	√ √
WiFi Update	WiFi EAP-TLS authentication protocol	-
Cybersecurity Features	√ ·	V
Security – User Accounts	User Accounts \\local user authentication create multiple local user accounts. And able to login/logout of the system	-
Table Side Remote Control Joy Stick Improvements	Updated skin layout to include Freeze, Camera and Measure.	V
Service Save logs update	system to provide a mechanism to transfer the below list of log files automatically to the SRS server	Save Log feature not automatic.
Security Hot-fix Installation	install a hotfix update via RUH (remote update handling) or USB drive.	install a hotfix update via RUH (remote update handling) or USB drive.
Study Back-up and Restore Improvements	Update: can restore settings from previous releases and not need to make the changes manually and can restore the same settings to multiple systems	Back-up and Restore Feature present
Motherboard version	MBM250	MBM201+ MBM250
Mechanical column upgrade	Unchanged	Lighter
Aluminum Card Cage	Unchanged	Weight reduction by 25 lbs
Monitor: 21" FPD	$\sqrt{}$	√
Output Display Standard (Track 3)	V	V
Patient Contact Materials	Tested to ISO 10993-1	Tested to ISO 10993-1
UL 60601-1 Certified	$\sqrt{}$	

Part 7. A brief discussion of nonclinical tests submitted, referenced, or relied on in the 510(k) for a determination of substantial equivalence.

The device has been evaluated for acoustic output, biocompatibility, cleaning and disinfection effectiveness as well as thermal, electrical, electromagnetic and mechanical safety and have been found to conform to applicable medical device safety standards.

The ACUSON SC2000 Ultrasound system complies with the following standards:

- AAMI / ANSI ES60601-1:2005/(R)2012 and A1:2012, C1:2009/(R)2012 and A2:2010/(R)2012, Medical electrical equipment Part 1: General requirements for basic safety and essential performance
 IEC 60601-1, Medical Electrical Equipment Part 1: General Requirements For Basic Safety And Essential Performance (IEC 60601-1:2005, MOD)
- IEC 60601-1-2: 2007(Third Edition), Medical electrical equipment Part 1-2: General requirements for basic safety and essential performance Collateral standard: Electromagnetic compatibility Requirements and tests
- IEC 60601-2-37:2007+A1:2015, Medical electrical equipment Part 2-37: Particular requirements for the basic safety and essential performance of ultrasound medical diagnostic and monitoring equipment
- IEC 60601-2-18:2009, Medical electrical equipment Part 2: Particular requirements for the safety of endoscopic equipment
- IEC 62304:2006 Medical Device Software Software Life Cycle Processes
- AIUM/NEMA UD-3:2004, Standard for Real Time Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic Ultrasound Equipment
- AIUM/NEMA UD-2:2004, Acoustic Output Measurement Standard for Diagnostic Ultrasound Equipment
- ISO 10993-1:2009, Biological evaluation of Biological evaluation of Medical Devices
- IEC 62366:2014, Medical Devices Application of Usability
- IEC 62359:2010, Ultrasonics Field characterization Test Methods for the determination of thermal and mechanical indices related to medical diagnostic ultrasonic fields.

Part 8. A summary discussion of the clinical tests submitted, referenced, or relied on for a determination of substantial equivalence.

Since the ACUSON SC2000 VC10A (v6.0) Diagnostic Ultrasound System uses the identical technology and principles of use as the existing predicate devices SC2000, VB21A (v5.1) K181098, clinical studies were not required to support substantial equivalence.

Part 9. Summary

Intended uses and other key features are consistent with traditional clinical practice and FDA guidelines. The design and development process of the manufacturer conforms to 21 CFR 820 Quality System Regulation and ISO 13485:2016 quality system standards. The product is designed to conform to applicable medical device safety standards and compliance is verified through independent evaluation with ongoing factory surveillance. The SC2000 Diagnostic ultrasound systems have accumulated a long history of safe and effective performance.

Therefore, it is the opinion of Siemens Medical Solutions USA, Inc. that the ACUSON SC2000 VC10A (v6.0) system is substantially equivalent with respect to safety and effectiveness to the currently cleared predicate devices for the U.S. market.